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L1 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN

RN 100-21-0 REGISTRY

ED Entered STN: 16 Nov 1984

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Terephthalic acid (7CI, 8CI)

OTHER NAMES:

CN 1,4-Dicarboxybenzene

CN 4-Carboxybenzoic acid

CN NSC 36973

CN p-Benzenedicarboxylic acid

CN p-Carboxybenzoic acid

CN p-Dicarboxybenzene

CN p-Phthalic acid

CN TPA

CN WR 16262

FS 3D CONCORD

DR 211863-90-0, 211863-92-2

MF C8 H6 O4

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*, DIOGENES, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, GMELIN\*, HODOC\*, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, NAPRALERT, NIOSHTIC, PDLCOM\*, PIRA, PROMT, PS, RTECS\*, SPECINFO, SYNTHLINE, TOXCENTER, TULSA, ULIDAT, USPAT2, USPATFULL, VTB (\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

9146 REFERENCES IN FILE CA (1907 TO DATE)
1861 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
9158 REFERENCES IN FILE CAPLUS (1907 TO DATE)
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> file caplus COST IN U.S. DOLLARS

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 100-21-0/prep9158 100-21-0 3321634 PREP/RL L2 2804 100-21-0/PREP (100-21-0 (L) PREP/RL) => s 100-21-0/proc9158 100-21-0 3703551 PROC/RL 700 100-21-0/PROC L3 '(100-21-0 (L) PROC/RL) => s 100-21-0/pur9158 100-21-0 213897 PUR/RL 476 100-21-0/PUR L4(100-21-0 (L) PUR/RL) => s 12 or 13 or 14 3336 L2 OR L3 OR L4 => s 15 and br and ni and mn 231380 BR 587562 NI 403130 MN 17 L5 AND BR AND NI AND MN L6 => s 16 and py<2004 23619165 PY<2004

17 L6 AND PY<2004

=> d 1-17 ibib abs hitstr

L7 ANSWER 1 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:845607 CAPLUS

DOCUMENT NUMBER:

137:338386

TITLE:

ь7

Method and catalyst system for preparing aromatic carboxylic acids from alkylaromatics by liquid-phase

oxidation

INVENTOR(S):

Park, Sang-Eon; Yoo, Jin S.; Jun, Ki-Won; Raju, David

B.; Kim, Young-Ho

PATENT ASSIGNEE(S):

Korea Institute of Chemical Technology, S. Korea

SOURCE:

U.S., 6 pp.

CODEN: USXXAM

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	 DATE
US 6476257	B1	20021105	US 2001-964750	 20010928 <
us 2002193631	A1	20021219		
KR 2002076879	А	20021011	KR 2001-17072	20010331 <
JP 2002332255	A2	20021122	JP 2002-94095	20020329 <
PRIORITY APPLN. INFO.:			KR 2001-17072	 20010331

AB In the title process, aromatic carboxylic acids (e.g., terephthalic acid) are prepared from alkylarom. hydrocarbons (e.g., p-xylene) by oxidation in an acetic acid solvent with oxygen-containing gas in the presence of a cobalt-manganese-bromine complex catalyst, to which nickel (e.g., nickel acetate) and carbon dioxide in appropriate amts. are added to increase catalyst activity. Nickel has a synergistic effect with carbon dioxide and maximizes the formation of the desired acid having the corresponding number of carboxylic groups to the number of alkyl groups in the reactant.

IT 100-21-0P, Terephthalic acid, preparation

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP

(Preparation)

(method and catalyst system with Co and Mn and Br and Ni and CO2 in acetic acid for preparing aromatic carboxylic acids from alkylaroms. by liquid-phase oxidation)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 2 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:825634 CAPLUS

DOCUMENT NUMBER:

138:287989

TITLE:

A study on process parameters in liquid phase

oxidation of p-xylene under the Co/Mn/

Br-based catalyst systems

AUTHOR(S):

Son, Young-Bae; Park, Kyung-Lyne; Yu, Tae-kong; Kim, Jung-Hi; Oh, In-Seok; Kim, Young-Ho; Yang, Hyun-Soo

CORPORATE SOURCE:

Dep. Fine Chemicals Eng. Chem., Chungnam National

Univ., S. Korea

SOURCE:

Nonmunjip - Ch'ungnam Taehakkyo Sanop Kisul Yon'guso (

2001), 16(2), 116-124

CODEN: NCTYEO

PUBLISHER:

Ch'ungnam Taehakkyo Sanop Kisul Yon'guso

DOCUMENT TYPE:

Journal

LANGUAGE:

Korean

AB Liquid phase oxidation of p-xylene using mol. oxygen has been carried out on the Co/Mn/Br-based catalyst system in acetic acid as a solvent. Process parameters(total pressure, partial pressure of oxygen and temperature) and kinetics of the reaction were studied as a basic study in the viewpoint for the development of new catalyst system and process. The total pressure above 20 atm and the partial pressure of oxygen above 5 atm

were required to keep the reactant in the liquid phase and to overcome the mass transfer limitation of oxygen. The oxidation rate and conversion were increased with increasing reaction temperature to the 100-190°C range. For a given initial concentration of p-xylene, the oxidation rate and the

For a given initial concentration of p-xylene, the oxidation rate and the apparent

activation energy were found to be of first-order and 4.24 kcal/mol, resp.

The effect of the third components added to the Co/Mn/Br

catalyst was also studied. It was found that Ni is only a good

additive that can promote the catalyst performance.

IT 100-21-0P, Terephthalic acid, preparation

RL: SPN (Synthetic preparation); PREP (Preparation)

(product; study, on process parameters in liquid phase oxidation of p-xylene under Co/Mn/Br-based catalyst systems)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 3 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:543886 CAPLUS

DOCUMENT NUMBER:

138:122875

TITLE:

Combined Promotional Effect of CO2 and Ni on

Co/Mn/Br Catalyst in the

Liquid-Phase Oxidation of p-Xylene

AUTHOR(S):

Raju Burri, David; Jun, Ki-Won; Yoo, Jin S.; Lee, Chul

Wee; Park, Sang-Eon

CORPORATE SOURCE:

Catalysis Center for Molecular Engineering, Korea Research Institute of Chemical Technology, Yuseong,

Daejeon, 305-600, S. Korea

SOURCE:

Catalysis Letters (2002), 81(3-4), 169-173

CODEN: CALEER; ISSN: 1011-372X Kluwer Academic/Plenum Publishers

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The promotional effects of CO2 and Ni were studied on Co/
Mn/Br catalyst in the liquid-phase oxidation of p-xylene to
terephthalic acid using mol. oxygen as an oxidant and acetic acid as a
solvent individually as well as in combination. The enhanced activity of
Co/Mn/Br catalyst was observed on both CO2 and Ni
promoters independently and also in combination. The activity enhancement
in the combination of CO2 and Ni promoters on this catalyst is
found to be remarkable.

IT 100-21-0P, Terephthalic acid, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (combined promotional effect of CO2 and Ni on Co/Mn/Br catalyst in liquid-phase oxidation of p-xylene)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

ANSWER 4 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

2000:619623 CAPLUS ACCESSION NUMBER:

134:42462 DOCUMENT NUMBER:

Formation and role of cobalt and manganese cluster TITLE:

complexes in the oxidation of p-xylene

Chavan, S. A.; Halligudi, S. B.; Srinivas, D.; AUTHOR(S):

Ratnasamy, P.

National Chemical Laboratory, Pune, 411008, India CORPORATE SOURCE:

Journal of Molecular Catalysis A: Chemical ( SOURCE:

2000), 161(1-2), 49-64

CODEN: JMCCF2; ISSN: 1381-1169

Elsevier Science B.V. PUBLISHER:

Journal DOCUMENT TYPE: English LANGUAGE:

The mono- and multinuclear metal complexes present during the aerial oxidation of p-xylene by the homogeneous catalyst systems viz. Co/Br

, Mn/Br, Co/Mn/Br, Co/Ce/ Br, Co/Zr/Br, Co/Mn/Zr/Br, Co/

Mn/Ce/Br, Ni/Mn/Br and

Ni/Mn/Zr/Br in acetic acid solvent have been

investigated by electronic and EPR spectroscopies. The reaction mixts.

contain, in addition to Co(OAc)2.4H2O and Mn (OAc) 2.4H2O, species like Co(OAc) Br, Co(OAc) 3,

Co3(O)(OAc)x, Mn3(O)(OAc)x and hetero-multinuclear complexes like

Co2Mn(O)(OAc)x and CoMn2(O)(OAc)x. While mononuclear Co(OAc)2, Mn

(OAc)2 and Co(OAc)Br complexes predominate in the initial stages

of the oxidation reaction, significant concns. of multinuclear Co(III) and

Mn(III) complexes are detected in the later stages. Zr(IV), when

present, facilitates the oxidation of Mn(II) to Mn(III),

a crucial step in the kinetic pathway of the oxidation reaction. EPR results indicate the presence of homo-nuclear Mn3(0)(OAc)x type clusters as the

major species with a minor component of CoMn2(0)(OAc)x complex in catalyst

systems with  $Co:Mn = 1:3 \pmod{1}$ . When an excess of cobalt catalyst is used (e.g., Co:Mn = 3:1) or when Zr and Br

are also present in optimal concns., the heteronuclear cluster

CoMn(O)(OAc) predominates while Co(O)(OAc) and CoMn(O)(OAc) occur as minor constituents. The yield of terephthalic acid (TA) is enhanced at high concns. of cluster complexes like Co2Mn(O)(OAc)x and CoMn2(O)(OAc)x. The

combination Co/Mn/Zr (3:1:0.1 mol), exhibits a high catalytic

activity and selectivity for terephthalic acid, especially in the presence of

optimal concentration of the bromide ion.

100-21-0P, Terephthalic acid, preparation IT

RL: SPN (Synthetic preparation); PREP (Preparation)

(formation and role of cobalt and manganese cluster complexes in oxidation of p-xylene to terephthalic acid)

100-21-0 CAPLUS RN

an

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME) CN

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS 39 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L7 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:130100 CAPLUS

DOCUMENT NUMBER: 126:131886

TITLE: Continuous process for the manufacture of terephthalic

acid of monomer purity by liquid-phase catalytic

oxidation of p-xylene

INVENTOR(S): Vovk, Ludmila Stepanovna; Linnik, Vladimir Petrovich;

Petrov, Viktor Pavlovich; Nazimok, Vladimir Filippovich; Tishchenko, Valery Ivanovich

PATENT ASSIGNEE(S): Mogilev Order of Lenin Proizvodstvennoe Obiedinenie

"Khimvolokno", Belarus

SOURCE: PCT Int. Appl., 23 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent Russian

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9641791	A1	19961227	WO 1996-BY4	19960508 <

W: JP, KR, RU, US

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE PRIORITY APPLN. INFO.: BY 1995-287 A 19950608

OTHER SOURCE(S): MARPAT 126:131886

AB The quality of terephthalic acid (I) is improved and unit consumption of AcOH (solvent) reduced by a combination of 2 interdependent processes for oxidation and recrystn. in a single aqueous AcOH solvent. The oxidation process is

carried out in a cascade of 2 reactors arranged in series at temps. of 180-199°C in the presence of a catalyst comprising mixts. of Co, Mn, Ni, Na, Li and K salts and mixts. of HBr and alkali metal bromides or brominated alkanes, e.g., MeCHBrCH2Br, as sources of Br. Intermediate products are removed from I at the stage of

recrystn. of I in AcOH under exptl. established conditions. The mother liquors are recirculated once the filtrates containing the raw (1st filtrate) and the pure (2nd filtrate) I have been separated during the 1st and 2nd oxidation stages, resp. A flow diagram of the process is included.

IT 100-21-0P, Terephthalic acid, preparation

RL: IMF (Industrial manufacture); PREP (Preparation)

(continuous process for the manufacture of terephthalic acid of monomer purity by liquid-phase catalytic oxidation of p-xylene)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 6 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:598540 CAPLUS

DOCUMENT NUMBER: 125:222740

TITLE: Manufacture and purification of terephthalic acid with

prevention of reactor corrosion

INVENTOR(S): Hara, Toshitsuna; Kasai, Yoshinori; Ishimaru,

Masaharu; Fukuda, Nobuo

PATENT ASSIGNEE(S): Mitsubishi Chemical Corp., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND \_\_\_\_\_\_ \_\_\_\_ JP 08193048 A2 19960730 JP 1995-5087 19950117 <--PRIORITY APPLN. INFO.: JP 1995-5087 19950117

The process comprises (1) oxidation of p-xylene in Br- and heavy metal-containing AcOH, (2) purification by hydrogenation of aqueous terephthalic acid

(I) with Pt-group catalysts, (3) crystallization of the purified I, and (4)

of wet I crystals in driers having Ni alloy parts in contact with the crystal heated at ≥100°. Oxidation of p-xylene in air in AcOH containing Co, Mn, and Br catalysts gave I, the aqueous I was fed into a distilling column containing Pd/C with H at 290°, crystallized by cooling to 150°, then dried in a dryer having Hastelloy C-lined parts (in contact with the crystal at 110°). The dryer had no corrosion over 1 yr.

100-21-0P, Terephthalic acid, preparation TT

RL: IMF (Industrial manufacture); PUR (Purification or recovery)

; PREP (Preparation)

(manufacture and purification of terephthalic acid with reactor corrosion prevention by Ni alloy linings)

100-21-0 CAPLUS RN

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME) CN

ANSWER 7 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1995:606805 CAPLUS

DOCUMENT NUMBER:

123:9155

TITLE:

Production method of high purity isomers of

benzenedicarboxylic acids

INVENTOR(S):

Nazimok, Vladimir Filippovich; Goncharova, Nadezhada Nikolaevna; Yurjev, Valerij Petrovich; Manzurov,

Vladimir Dmitrievich

PATENT ASSIGNEE(S):

Samsung General Chemicals Co., Ltd., S. Korea;

Joint-Stock Company of Research and Design Institute

of Monomers

SOURCE:

PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT	NO.			KIN	D	DATE			APPL	CAT	ION 1	.00		DA	ATE		
						_												
WO	9509	143			A1		1995	0406	1	WO 1	993-	KR10	6		19	9931	130 <	:
	W:	AT,	AU,	BB,	BG,	BR,	CA,	CH,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,	JP,	ΚZ,	
							NO,											
	RW:	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG			·

RU	2047594	C1	19951110	RU	1993-46190		19930928	<
RU	2047595	C1	19951110	RU	1993-46191		19930928	<
KR	9700136	B1	19970104	KR	1993-21276		19931014	
	9305996	Α	19971021	BR	1993-5996		19931120	<
	2128719	AA	19950329	CA	1993-2128719		19931130	<
CA	2128719	С	19980609					
AU	9455763	A1	19950418	ΑU	1994-55763		19931130	
GB	2286588	A1	19950823	GB	1994-15915		19931130	<
	2286588	B2	19960911					
ES	2081265	A1	19960216	ES	1994-50020		19931130	<
ES	2081265	B1	19961016					
DE	4397599	T	19970724	DE	1993-4397599		19931130	<
DE	4397599	C2	19980219					
RO	113850	B1	19981130		1994-1218	•	19931130	
$\mathtt{PL}$	175685	B1	19990129	PL	1993-308537		19931130	
JP	3009223	B2	20000214	JP	1994-516877		19931130	<
JР	08506571	T2	19960716					
SK	280582	В6	20000410	SK	1994-877		19931130	
CN	1103860	Α	19950621	CN	1994-100679		19940112	<
CN	1050118	В	20000308		•			
BE	1008546	A4	19960604		1994-702		19940726	
FR	2710638	A1	19950407	FR	1994-11253		19940921	<
FR	2710638	B1	19960426					
PRIORIT	Y APPLN. INFO.:				1993-46190	Α	19930928	
					1993-46191	Α	19930928	
				WO	1993-KR106	W	19931130	

OTHER SOURCE(S): CASREACT 123:9155

AB An improved process for producing highly purified benzenedicarboxylic acid isomers, having color index of ≤10°H and containing <0.0025% carboxybenzyldehyde (CBA)impurity, without an addnl. catalytic reductive purification step, comprises (a) an oxidation step wherein xylene isomer is oxidized with mol. O or mol. O containing gas in the presence of a catalyst system composed of Co Mn, Br and at least one selected from Ni, Cr, Zr and Ce in low aliphatic carboxylic acid; and (b) an

extraction/post-oxidation step wherein the oxidation product is crystalline to give a cake

of crude benzenedicarboxylic acid isomer, the cake is reslurried by adding lower aliphatic carboxylic acid solvent thereto followed by heating in order to extract impurities contained therein into the solvent, and the resulting slurry is oxidized with said catalyst system at a temperature of 2-80° lower than that of said heating, each of said oxidation and

extraction/post-oxidation

being carried out once or twice, provided that any one or both of said steps should be carried out twice. According to the invention, the solvent employed to extract impurities is recycled from the subsequent sidation

steps. P-xylene, AcOH, H2O, Co, Mn, Ni and Br were heated to 160°, oxidation effected at 198° and 19 kg/cm2 for 40 min resulting in 20% terephthalic acid, 25 ppm 4-carboxybenzaldehyde and color index of 8°H.

1T 100-21-0P, 1,4-Benzenedicarboxylic acid, preparation
RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
(Preparation)

(production method of high purity isomers of benzenedicarboxylic acids)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1992:129688 CAPLUS

DOCUMENT NUMBER: 116:129688

TITLE: Developments in terephthalic acid manufacture

AUTHOR(S): Nazimok, V. F.; Pivovar, L. M.

CORPORATE SOURCE: USSF

SOURCE: Khimicheskie Volokna (1991), (6), 38-40

CODEN: KVLKA4; ISSN: 0023-1118

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB A scheme is presented for the continuous manufacture of terephthalic acid (I) by liquid-phase oxidation of p-xylene (II) in HOAc in the presence of a Co-

Mn-Ni-Br catalyst. The oxidation is conducted in

3 steps. High-quality I can be obtained at 180-200° with 17% II in

the starting mixture on using an optimized catalyst composition Aliphatic

C3-6-hydrocarbons are used as promoters.

IT 100-21-0P, Terephthalic acid, preparation

RL: IMF (Industrial manufacture); PREP (Preparation)

(manufacture of, by liquid-phase oxidn.of xylene, technol. for continuous)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1990:197849 CAPLUS

DOCUMENT NUMBER: 112:197849

TITLE: Continuous two-stage oxidation of aromatic

hydrocarbons to aromatic carboxylic acids in an

aqueous system

INVENTOR(S): Nowicki, Neal R.; Lowry, James D., Jr.

PATENT ASSIGNEE(S): Amoco Corp., USA

SOURCE: U.S., 5 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
US 4892970 PRIORITY APPLN. INFO.:	Α	19900109	US 1985-814510 US 1985-814510	19851230 < 19851230	

AB Benzenes (I) disubstituted with oxidizable substituents such as (hydroxy)alkyl, aldehyde, or carboalkyl are oxidized to their corresponding carboxylic acid derivs. in an aqueous solvent system in a continuous, 2-stage process comprising (1) partial (75-85%) oxidation of a feed stock containing I 100, H2O .apprx.5-100, ionic catalytic metal (Mn and at least one of Co, Ni, Zr or its mixture) 0.1-5.0 weight parts, and Br .apprx.10-300 atom % (based on the total catalytic metal) with a stoichiometric excess of an O-containing gas at .apprx.300-410°F and a pressure high enough to maintain a liquid phase, wherein byproduct formation is minimized and (2) addition of supplement Br to the first stage reaction mixture to raise the

concentration of Br (50-500 atom % based on the metal) and of an O-containing gas at .apprx.400-480°F, wherein the added Br effectively suppresses decarboxylation reactions that typically occur at higher oxidation temperature Thus, p-xylene (II) was oxidized to terephthalic acid

(III) in H2O in the presence of Co, Mm, and Br in the first stage at 380°F and 525 psig (8 psi 0) using Co and Mm 2.7 weight%), Mm/Co (3.0 % of II), Br/Mm + Co (0.35), H2O/II (0.3) and in the second stage at 453°F and 510 psig (11 psi 0) using Co and Mm (3.3% of II), Br/Mm + Co (2.8, 8 fold increase) and H2O/II (2.3). The slurry obtained contained toluic acid 0.08, 4-(HO2C)C6H4CHO 0.18, III 33.0, and high mol. weight component 0.20 weight %.

100-21-0P, 1,4-Benzenedicarboxylic acid, preparation RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by oxidation of p-xylene)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 10 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1989:192438 CAPLUS

DOCUMENT NUMBER: 110:192438

TITLE: Oxidation process for the manufacture of aromatic

acids from alkylaromatic compounds

INVENTOR(S): Partenheimer, Walter; Schammel, Wayne P.

PATENT ASSIGNEE(S): Amoco Corp., USA

SOURCE: U.S., 5 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4786753	A	19881122	US 1987-50860	19870518 <
EP 362443	A1	19900411	EP 1988-309255	19881005 <
EP 362443	B1	19940112		
R: BE, DE, ES,	FR, GB	, IT, NL		
ES 2047557	Т3	19940301	ES 1988-309255	19881005 <
JP 02138149	A2	19900528	JP 1988-263044	19881020 <
JP 2746346	B2	19980506		
CN 1042534	A	19900530	CN 1988-107608	19881105 <
CN 1024660	В	19940525		
PRIORITY APPLN. INFO.:			US 1987-50860	19870518
			EP 1988-309255	A 19881005

OTHER SOURCE(S): CASREACT 110:192438

AB A process of oxidizing di- and trimethylbenzenes with O2 to benzenedi- and -tricarboxylic acids in the liquid phase in the presence of a C<5 aliphatic acid, H2O, or a mixture of the aliphatic acid and H2O at .apprx.100-260° and at a pressure to maintain as liquid phase 70-80% of the reaction medium comprised conducting said oxidation in the presence of a catalyst system comprising a source of Br with Ni, Zr, and Mn wherein for each g-mol of p-xylene, m-xylene, or pseudocumene in the

oxidation there is from .apprx.4 to .apprx.20 mg-atom Ni, .apprx.0.10 to .apprx.0.30 mg-atom Zr, .apprx.2 to .apprx.10 mg-atom total Mn, and from .apprx.8 to 24 mg-atom Br. The addition of Ni and Zr to oxidation catalysis provided by heavy, transition metal-Br ion combination containing at least Mn ion uniquely increases catalytic activity of said combination for converting Me groups to CO2H on the benzene nucleus and enables the elimination of the expensive catalyst Co. The preparation of phthalic acid, isophthalic acid, and trimellitic acid from p-Me2C6H4, m-Me2C6H4, and pseudocumene, resp., using various Ni, Zr, Mn, and Br combinations is given.

100-21-0P, Terephthalic acid, preparation IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by catalytic oxidation of p-xylene)

RN 100-21-0 CAPLUS

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME) CN

CAPLUS COPYRIGHT 2005 ACS on STN ANSWER 11 OF 17

ACCESSION NUMBER: 1986:68602 CAPLUS

DOCUMENT NUMBER:

104:68602

TITLE:

Tere- or isophthalic acids

INVENTOR(S):

Nazimok, V. F.; Golubev, G. S.; Boyarkin, M. A.;

Manzurov, V. D.; Yur'ev, V. P.; Khomin, V. V.;

Pakhorukov, V. A.

PATENT ASSIGNEE(S):

All-Union Scientific-Research and Design Institute of

Monomers, USSR

SOURCE:

U.S.S.R. From: Otkrytiya, Izobret. 1985, (29), 102.

CODEN: URXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Russian

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
SU 1171452	A1	19850807	SU 1983-3620406	19830510 <
PRIORITY APPLN. INFO.:			SU 1983-3620406	19830510
AB Tere- and isophtha	lic acio	ds are prepa	red by liquid-phase o	xidation of p- or
m-xylene, resp., w	ith a ga	as containin	g mol. O in HOAc; in	the 1st stage at
200-215°/20-26 atm				
Ni salts and Br co	mpds., v	with subsequ	ent treatment of	
the reaction mater	ial, in	the 2nd sta	ge, at 180-200°/8.2-1	.0 atm
gage with steam an	d gas m	ixture The	reaction mixture was	further treated with
0.3-0.6 parts by w	eight o:	f a reflux s	colution from crystall	ization zones,
containing HOAc				
80-88, H2O 10-16.	and ROR	1 (R, R1 = H)	(, Me, MeO) 1.5-4.0%.	

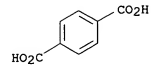
8U-88, HZO LU-L6, and RORL (R, RL

IT 100-21-0P, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by oxidation of xylene)

100-21-0 CAPLUS RN

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME) CN



L7 ANSWER 12 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1980:567949 CAPLUS

DOCUMENT NUMBER: 93:167949

TITLE: Method of preparing terephthalic acid

INVENTOR(S): Nazimok, V. F.; Kulakov, V. N.; Manzurov, V. D.;
Boyarkin, M. A.; Golubev, G. S.; Simonova, T. A.;

Valieva, R. A.; Petrov, A. A.; Zernov, P. N.; et al. All-Union Scientific-Research and Design Institute of

PATENT ASSIGNEE(S): All-Union Scientific-Research Monomers, USSR

SOURCE: U.S.S.R. From: Otkrytiya, Izobret., Prom. Obraztsy,

Tovarnye Znaki 1980, (21), 135.

CODEN: URXXAF

DOCUMENT TYPE: Patent LANGUAGE: Russian

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
SU 739062 PRIORITY APPLN. INFO.: AB The title compound	T was pre		SU 1978-2591177 SU 1978-2591177 A liquid phase oxidation	19780316 < 19780316 of p-xylene in

in presence of a Co, Mn, Ni catalyst and Brinitiator. The process was carried out in two stages. The first stage
was at 210-30° and 24-7 atms in presence of a catalyst with total
concentration of Co and Mn 0.045-0.15% of the reaction mixture and weight
concentration of Ni 0.0005-0.005% and ratio of Mn:Ni
= 30-100. The second stage was at 160-210° and 5-10 atms. The
mixture contained 40-70% AcOH, 10-20% H2O, 15-50% N and 0.1-1.5% O.

IT 100-21-0P, preparation

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by oxidation of xylene)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 13 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1980:446199 CAPLUS

DOCUMENT NUMBER: 93:46199

TITLE: Recovery of nondissociated bromine in spent oxidation

catalyst solution after terephthalic acid production

INVENTOR(S): Takeuchi, Hiroshi; Saeki, Satoshi

PATENT ASSIGNEE(S): Asahi Chemical Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
		A2	19800226	JP 1978-99101	
PRIC	ORITY APPLN. INFO.:				19780816
AB	p-Xylene is oxidize	ed with	O in AcOH c	ontaining heavy-metal b	romide, and the
	mother liquor is da	istilled	d first to re	ecover AcOH and then at	P mm Hg pressure
				9.57 - 3330/(t + 348).	
	Thus, AcOH 20 kg co	ontainir	ng CoBr2.6H2	0 20 and Mn (OAc) 2.4H20	20 g in
				nd 20 kg/cm2 gage with	
	10% p-xylene and 5	kg/h ai	ir for 5 h,	the product was collect	ed
	continuously in a	crystall	lization ves	sel at $180^{\circ}$ and $10$ kg/c	m2 gage and
				taining some p-(HO2C)2C	6H4 (I), H2O 5%, Co
	750, Mn 700, Fe 52,				
	and undissocd. Br	1800 ppr	n was fed at	8 kg/h at the top of a	
	distillation colum	n filled	d with Rasch	ig rings at $114^\circ$ to sep	. AcOH and
				om was fed 1 kg/h into	an Arther Smith
•	thin-film evaporate				
				ycled along with Co and	
		I yield	d containing	190-210 ppm 4-HCOC6H6C	OZH was
	96-8% in 6 cycles.		•		
TΨ	100-21-00 preparat	tion		•	•

100-21-0P, preparation IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by oxidation of p-xylene)

100-21-0 CAPLUS RN

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME) CN

CAPLUS COPYRIGHT 2005 ACS on STN ANSWER 14 OF 17

ACCESSION NUMBER:

1979:204824 CAPLUS 90:204824

DOCUMENT NUMBER: TITLE:

Recovery of liquid-phase oxidation catalyst and

solvent

INVENTOR(S):

Miyake, Tetsuya; Takeuchi, Hiroshi; Tauchi, Masatoshi;

Saeki, Miciyuki; Saka, Kazuki

PATENT ASSIGNEE(S):

Asahi Chemical Industry Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 53104590	A2	19780911	JP 1977-19667	19770224 <
PRIORITY APPLN. INFO.:		•	JP 1977-19667 A	
			6-42-3] with mol. O to	
			talysts containing Co,	
<b>Br</b> in lower alipha	tic mond	carboxylic a	acid, the reaction mixt	ure after

Co and Mn ions and Cu and Zn ions if present, and the metal compds. were recovered and recycled. The effluent from the anion

separation of II was treated with Br--type anion exchanger to remove

exchanger treatment was treated with a carboxylate-type, weakly basic anion exchanger at 20-150° to remove Br ion and Ni if present, which were also recovered and recycled. For example, a mixture of 20 kg AcOH [64-19-7], 20 g (as Co) CoBr2.6H2O and 20 g (as Mn) Mn (OAc) 2.4H2O at 200 $^{\circ}$ /20 kg/cm2 gage was fed with 10% I solution in AcOH at the rate of 20 g/h and 20 kg/h air, maintaining the reaction mixture at 180°/10 kg/cm2 gage for 5 h, and the reaction mixture was continuously discharged, separated from II, and passed through a pyridinium bromide-type anion exchanger to give an effluent (III) containing Co <1, Mn <1, and Br 2400 ppm, compared with 920, 910, and 2700, resp., for the reaction mixture The spent resin was eluted with AcOH containing 2% water to give a 30 mL solution containing Co 15,000, Mn 14,500, and Br 4200 ppm. III passed through a pyridinium acetate-type anion-exchange column at 80° to give an effluent containing <10 ppm Br; elution of the resin with AcOH gave a solution containing 40,000 ppm Br. AcOH was recovered in 93.8% yield by fractional distillation

IT 100-21-0P, preparation

RL: IMF (Industrial manufacture); PREP (Preparation) (manufacture of, from xylene, recovery and recycle of solvent and oxidation catalysts in)

100-21-0 CAPLUS RN

1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

CAPLUS COPYRIGHT 2005 ACS on STN ANSWER 15 OF 17

ACCESSION NUMBER:

1978:604884 CAPLUS

DOCUMENT NUMBER:

89:204884

TITLE:

Recovery of cobalt-manganese-bromide catalyst

INVENTOR(S):

Miyake, Tetsuya; Takeuchi, Hiroshi; Tauchi, Masatoshi;

Saeki, Michiyuki; Saka, Kazuki

PATENT ASSIGNEE(S):

Asahi Chemical Industry Co., Ltd., Japan

SOURCE:

CN

Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 53102290	A2	19780906	JP 1977-16756	19770218 <
DTODITY APPIN INFO :			JP 1977-16756 A	19//0210

With Co and Mn catalyst and Br as accelerator, alkylbenzene or its derivative is oxidized with 0 in lower fatty acid solvent into the corresponding carboxylic acid, the resulting solution is contacted with anion-exchange resin containing pyridine ring in the bromide or solvent anion form, and eluted to recover Co and Mn with or without Br. Thus, p-xylene was oxidized in AcOH containing Co, Mn, and Br in a 2 L Ti vessel at 205° and 15 atm, cooled to 100°, and filtered to sep. terephthalic acid; the 500 mL solution containing p-xylene 50, 4-carboxybenzaldehyde 450, p-cresol 20 ppm, Co 0.10, Mn 0.02, Br 0.20, Fe 0.005, Cr 0.003, and NO 0.002% was passed through a pyridine anion exchanger column 12 diameter + 100 mm at 80° and 200 mL/h. The effluent contained Co, Mn

<0.001 each, **Br** <0.002, Fe 0.0049, Cr 0.0028, and **Ni** 0.0017%, vs. 0.072, 0.012, 0.033, 0.0047, 0.0029, and 0.0019% with Dowex 1X4.

IT 100-21-0P, preparation

RL: PREP (Preparation)

(manufacture of, by oxidation of xylene, recovery of cobalt-manganese catalysts

for)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 16 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1977:106180 CAPLUS

DOCUMENT NUMBER: 86:106180

TITLE: High-purity terephthalic acid INVENTOR(S): Namie, Koushi; Takeda, Shinichi

PATENT ASSIGNEE(S): Teijin Hercules Chemical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
	JP 51108028	A2	19760925	JP 1975-31829		19750318	<
	JP 59008253	B4	19840223	•			
PRIC	DRITY APPLN. INFO.:			JP 1975-31829			
AB	carboxylic acid sol	lvent wa cration	is carried of in the sol	p-xylene to terephtha out at 170-280° with vent and ≥0.8 time	500-	6000	Oina
	weight of Br (as HE AcOH containing 0.4	3r) base 146 g Mr 3 1:1 Mr	ed on the to (OAc)2, 0. n-Ni in AcO	otal metal amount TI 424 g Ni(OAc)2, and H, Br/(	nus,	200 g	,
	<pre>Mn + Ni) = 3] was t L./min air at 220°, terephthalic acid o ppm, resp., with 0.</pre>	/20 kg/c containi	cm2 in an a ing 190 ppm	n p-xylene and 1.5 utoclave to give 97% p-carboxybenzaldehy	de, v	s. 92% an	d 1240

IT 100-21-0P, reactions

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, from p-xylene)

RN 100-21-0 CAPLUS

CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)

L7 ANSWER 17 OF 17 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1975:64116 CAPLUS

DOCUMENT NUMBER:

82:64116

TITLE:

Recovery of heavy metal bromides and hydrogen bromide

from reaction mixture for producing terephthalic acid

Shigeyasu, Mottoo; Ozaki, Takeo; Kusano, Nobuo

INVENTOR(S):
PATENT ASSIGNEE(S):

Matsuyama Petrochemicals Inc.

SOURCE:

Ger. Offen., 38 pp.

CODEN: GWXXBX

Patent

DOCUMENT TYPE: LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2415393	A1	19741024	DE 1974-2415393	•	19740329 <
DE 2415393 DE 2415393	C3 B2	19791018 19790301			10700000
JP 49123192 JP 55007300	A2 B4	19741125 19800223	JP 1973-36351		19730330 <
JP 54019399 JP 49133290	B4 A2	19790714 19741220	JP 1973-47826		19730426 <
BE 813080 PRIORITY APPLN. INFO.:	A1	19740715	BE 1974-142667 JP 1973-36351	A	19740329 < 19730330
			JP 1973-47826	Α	19730426

AB Br, Co, and Mn are recovered from the residue obtained after the removal of terephthalic acid (I) and the remaining solvent from the reaction mix for the preparation of I by oxidation of p-dialkylbenzene in the

presence of brominated heavy metal catalysts, particularly Co and Mn, by treating the residue with H2O and O2 in the presence of a sulfide, such as H2S, Na2S, NaHS, K2S, NH4HS, etc., separating precipitated sol.

impurities which may include Fe and Cr oxides and sulfides of Pb, Mo, Cu, Ni, or Cd, treating the aqueous solution with a strong acid cation exchange resin in H+ form, to absorb the Co and Mn ions, and distilling the remaining solution to recover HBr. The absorbed Co and Mn ions are removed from the ion exchange resin by eluting with aqueous HBr and the eluant distilled to recover excess HBr and isolate bromides of Co and Mn. Thus, to a tarry residue 700 recovered after the preparation of I with AcOH solvent, CoBr2, Mn acetate, and HBr and afer removal of I and AcOH, which residue contains Co 4.6, Mn 0.27, Fe 0.32, Cr 0.09, Br 4.5%, Cu 12, Mo 6, and Pb 4 ppm, H2O 1050 kg is added, heated to 70° for 2 hr with stirring and 300 l./hr of air together with 40 1. of H2S passed through, passed to a crystallization vessel where it is cooled to 30°, centrifuged to remove solids to provide 950 kg of filtrate containing Co 3, Mn 0.15, Br 3%, as well as Fe 59, Cr 5.4, Cu 1.5, Mo 0.7, and Pb 0.5 ppm. The yield of Co extracted is 89%. The Co and Mn ions are absorbed in an ion exchange resin column. When the resin has absorbed its capacity of metal ions, it is 1st washed with H2O (the resultant H2O being used for the next extraction step), then washed with 10% AcOH to remove organic impurities, and

then

ΙT

eluted with 3N HBr to desorb Co and Mn; the resultant eluant containing CoBr2, MnBr2, and HBr, is distilled recovering excess HBr and leaving

a product containing Co 23.8, Mn 1.2, Fe 0.04, Cr 0.004, Br 74.6%, Cu 3, Mo 3, and Pb 3 ppm. The yield of Co from the extract is 97%. 100-21-0P, preparation

RL: IMF (Industrial manufacture); PREP (Preparation)
(waste water from manufacture of, hydrogen bromide and metal bromide recovery from, ion exchange and distillation in)

RN 100-21-0 CAPLUS CN 1,4-Benzenedicarboxylic acid (9CI) (CA INDEX NAME)